## **WEST Search History**

DATE: Wednesday, July 30, 2003

Set Name side by side	Query	Hit Count	Set Name result set
DB = USPT, F	PGPB; PLUR=YES; OP=ADJ	-	
L2	L1 and env	41	L2
L1	HIV and Group O	481	L1

END OF SEARCH HISTORY

07nov01 15:15:53 User208669 Session D1928.1

\$0.26 0.073 DialUnits File1

\$0.26 Estimated cost File1

\$0.26 Estimated cost this search

\$0.26 Estimated total session cost 0.073 DialUnits

File 155:MEDLINE(R) 1966-2001/Dec W1

\*File 155: From 11/5/2001 the NLM will not update Medline until early 2002 This is the period in which the NLM completes the annual re-indexing

Set Items Description

? s hiv and o

SI 1316 HIV AND O

? s env and s1

5962 ENV

1316 SI

105 ENV AND S1 S2

? s py<1998

Processing

S3 9572908 PY<1998

105 S2 ? s s2 and s3

9572908 S3

49 S2 AND S3

DIALOG(R)File 155:MEDLINE(R)

Diversity of the immunodominant epitope of gp41 of HIV-1 subtype O and ts validity for antibody detection.

Eberle J, Loussert-Ajaka I, Brust S, Zekeng L, Hauser PH, Kaptue L, Knapp

S; Damond F; Saragosti S; Simon F; Gurtler LG

Pettenkofer Institute, University of Munchen, Germany.

Journal of virological methods (NETHERLANDS) Aug 1997, 67 (1) p85-91

ISSN 0166-0934 Journal Code: HQR

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

subtype O isolates, ANT70, MVP-5180 and VAU. All HIV-1 subtype O isolates from Cameroon, 11 from France and one from Germany were sequenced. The The immunodominant regions of the gp41 from 13 HIV-1 subtype O strains amino acid sequences were compared to those of the 3 published HIV-1

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A 25 residue peptide from the immunodominant domain of gp41 of the MVP-5180 (CM.6778 and CM.8161) showed an acidic amino acid in this loop. None of the representing the immunodominant epitope of the strain MVP-5180 successfully solates showed the same amino acid sequence in this immunodominant region. enzyme labeled anti-human IgG. Out of 111 anti-HIV-1 positive specimens, collected mainly from Cameroonian HIV infected patients, only 10 were not reactive in this assay. The 42 anti-HIV-1 subtype O positive specimens gave microtiter plates. Antibody binding was detected by indirect ELISA using an had a very conserved amino acid sequence in this region and showed a subtype O specific structure. Within the cysteine loop there was a positive infected patients, while some assays without HIV-O specific antigens charge of two basic amino acids, arginine and lysine. Only two strains of adding such a peptide for correct identification of HIV-1 subtype O detected all the anti-HIV-O sera so far tested, pointing to the importance strain was synthesized, cycled to form the cysteine-loop and coated to all a reaction above cut off. Despite the diversity found in the amino acid sequences within the 25 isolates a peptide-based indirect ELISA partially fail to detect all anti-HIV-O specimens.

Record Date Created: 19971014

DIALOG(R)File 155:MEDLINE(R)

Envelope sequence variability and serologic characterization of HIV type group O isolates from equatorial guinea.

Hunt JC; Golden AM; Lund JK; Gurtler LG; Zekeng L; Obiang J; Kaptue L;

Hampl H; Vallari A; Devare SG

AIDS Research and Retrovirus Discovery, Abbott Laboratories, North

Chicago, Illinois 60064, USA.

AIDS research and human retroviruses (UNITED STATES) Aug 10 1997, 13

(12) p995-1005, ISSN 0889-2229 Journal Code: ART

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

were either RIGPLAWY (one isolate), RIGPMAWY (two isolates), or GLGPLAVY and Western blot. Degenerate primers, designed from HIV-1 group O published EG sera clustered more closely with HIV Env sequences of group O compared sequences, were used to PCR amplify envelope (env) gene sequences from the suspect EG sera. A complete envelope gene sequence from each serum was determined from the overlapping env gene fragments. Analysis (PHYLIP (one isolate). The V3 tip tetrameric sequence GPLA is represented only once nucleotide sequences) indicated that the amino acid sequences obtained from to group M. The amino acid sequences at the octameric tip of the V3 loop Four sera from Equatorial Guinea (EG) suspected to contain antibody package of programs) of Env amino acid sequences (translated from unusual and differential serologic reactivity in selected commercial assays against HIV-1 group O-related viruses were identified on the basis of

RLLALETLIQNQQLLNLWGCKGR(K)L(I)VCYTSVK(T)W, whereas sequence from the O-related EG samples. The gp41 immunodominant regions (IDR) protein in the 1995 HIV (Los Alamos) database, but was present in two of our group sequences were identical for sequences from three of the sera,

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group O isolate ANT70, VAU, or MVP5180. Antibody in each EG serum directed sequences from the ANT70 or MVP5180 IDRs, but were most reactive against derived from EG sera were unique compared to those reported for other HIV-1 comprising the IDR of HIV-1 group M (subtype B consensus) or HIV-2 serum contained three changes as noted in parentheses. IDR sequences the sequences derived from the samples themselves. Little or no serologic against the IDR could be detected using synthetic peptides comprising reactivity was detected when EG sera were reacted against peptides

Record Date Created: 19971010

DIALOG(R)File 155:MEDLINE(R)

Hunt JC; Brennan CA; Golden AM; Yamaguchi J; Lund JK; Vallari AS; Hickman Molecular analyses of HIV-1 group O and HIV-2 variants from Africa. RK; Zekeng L; Gurtler LG; Hampl H; Kaptue L; Devare SG

Abbott Laboratories, North Chicago, IL-60064, USA.

Leukemia (ENGLAND) Apr 1997, 11 Suppl 3 p138-41, ISSN 0887-6924

Iournal Code: LEU

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

sequences at the octameric tip of the V3 loop were RIGPLAWY, RIGPMAWY, or closely with HIVANT70 compared to other Group O sequences. The amino acid immuno-dominant region (IDR) sequences derived from EG sera were unique in GLGPLAVY. The tetrameric tip GPLA is represented only once in the published 1994 HIV database (Los Alamos) but was present in 2 of 4 of EG samples. The extracted from these samples was used to PCR amplify overlapping sequences that none of the sequences were completely homologous to other HIV-1 group Guinea (EG) led to identification of 4 HIV-1 Group O variants. Viral RNA, detection by serologic and genetic techniques. To characterize the sequence selected commercial and research assays. Analysis of sera from Equatorial indicated that the V3 loop nucleotide and protein sequences aligned more of the entire envelope gene using multiple primer pairs. Sequence analysis variation and its correlation to serologic diversity of HIV-1 Group O and O variants. Further, the HIV-1 group O sequence variation could be Genetic variation among HIV isolates creates challenges for their HIV-2 isolates, samples were identified by differential reactivity in correlated with differential serologic reactivity using IDR peptides.

WGCAFRQVCHT. This region is absolutely conserved among the majority of currently known HIV-2 and related SIV viruses (1994 HIV database). One subtype B sample had a unique sequence immediately adjacent to the IDR, serologic detection of HIV-2, plasma samples from Western Africa were to env proteins. PCR amplification and sequence analysis of the gag, env V3 evaluated. Eight samples were selected based on weak scrologic reactivity oop, and env IDR regions indicated that the samples could be classified as however, this did not change the serologic detection using a HIV-2 IDR HIV-1. To further establish a correlation between the genetic diversity and subtypes A (4 samples), B (3 samples) and D (1 sample). Across the subtypes, there was conservation in the IDR region of the sequence specific monoclonal antibody.

Record Date Created: 19970807

DIALOG(R)File 155:MEDLINE(R)

Env gene characterization of the first HIV type 1 group O Spanish solate.

Mas A; Quinones-Mateu E; Soriano V; Domingo E

Servicio de Enfermedades Infecciosas, Instituto de Salud Carlos III,

Madrid, Spain.

ADS research and human retroviruses (UNITED STATES) Nov 20 1996, 12 (17) p1647-9, ISSN 0889-2229 Journal Code: ART

Document type: Journal Article

Languages: ENGLISH

Record type: Completed

Record Date Created: 19970227

DIALOG(R)File 155:MEDLINE(R)

V3 loop sequence analysis of seven HIV type 1 group O isolates phenotyped in peripheral blood mononuclear cells and MT-2 cells. De Jong J; Simon F; Van der Groen G; Baan E; Saragosti S; Brun-Vezinet F;

Department of Human Retrovirology, Academic Medical Centre, Amsterdam, The Netherlands.

Goudsmit J

AIDS research and human retroviruses (UNITED STATES) Nov 1 1996, 12

(16) p1503-7, ISSN 0889-2229 Journal Code: ART

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

only non-syncytium-inducing (NSI) viruses. The syncytium-inducing capacity HIV-1-infected individuals from which syncytium-inducing (SI) viruses are isolated most often progress more rapidly to AIDS than individuals carrying of virus isolates is commonly determined in conjunction to replication in

Compared to HIV-1, the sequence information on HIV-2 isolates is relatively

imited, though the HIV-2 isolates also show genetic variation similar to

MT-2 cells. Comparison of HIV-1 env sequences and a site-directed mutagenesis study have indicated that the presence of a positively charged amino acid at position 11 or 25 in the V3 loop is minimally required for the SI capacity of HIV-1 subtype B viruses. Studies have also shown a similar correlation between positively charged signature amino acids in the V3 loop and syncytium formation in MT-2 cells for HIV-1 subtypes A, D, and E. In the present study virus phenotype was determined and compared to the V3 loop sequence of seven HIV-1 group O isolates. Three of the HIV-1 group O isolates showed the NSI/non-MT-2 tropic phenotype and two showed the SI/MT-2 tropic phenotype. The V3 loop of the two SI/MT-2 tropic isolates had a high net positive charge and contained a positively charged amino acid at position 11 or 25. The V3 loop of the two NSI/MT-2 tropic isolates had a low net positive charge and contained a single positively charged amino acid at position 37.

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Record Date Created: 19970224

4/7/14

DIALOG(R)File 155:MEDLINE(R)

An efficient method for the rescue and analysis of functional HIV-1 env genes: evidence for recombination in the vicinity of the tat/rev splice site.

Douglas NW; Knight AI; Hayhurst A; Barrett WY; Kevany MJ; Daniels RS Virology Division, National Institute of Medical Research, Mill Hill, London, UK.

AIDS (UNITED STATES) Jan 1996, 10 (1) p39-46, ISSN 0269-9370 fournal Code: AID

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

genes with open reading frames. METHODS: A nested polymerase chain reaction structure of gp160. RESULTS: From random patient samples collected in directly from patient samples. DESIGN: HIV exists as a quasispecies which cloned into a T7-promoter-containing vector. Expression of gp160 in CV-1 (PCR) approach has been used to rescue intact (2.6 kb) env genes, which are London clinics, only HIV-1 subtype B was found. Two of the samples Translation products are analysed in relation to the known immunogenic characterization of full-length expression-competent HIV-1 env genes can be disturbed by in vitro culture, in which numerous members of the dominant role in disease progression. Since env gene translation products OBJECTIVE: To establish a robust procedure for the isolation and play major roles in the initiation and spread of infection we need to study cells is detected by Western blot. Expression-competent clones are sequenced and resulting sequences used for phylogenetic studies. population are likely to be defective due to the high error rate of the viral reverse transcriptase. Defective viruses are unlikely to play a

contained viruses with an additional pair of cysteine residues in their V1 regions. For samples collected in Uganda, HIV-1 subtypes A, D and an A/D recombinant were recovered. CONCLUSION: An effective procedure is described for the isolation of HIV-1 env genes directly from patient samples, which has worked for A, B and D subtypes to date. The PCR primers can be utilized with other subtypes with the possible exception of subtype O viruses. Phylogenetic analyses revealed the potential importance of a G/C-rich region near the tat/rev splice site as a site of recombination. The sequences and translation products generated may be more relevant to disease progression in vivo and vaccine formulations than those obtained from viruses selected in long-term culture.

Record Date Created: 19961031

4/1/1

DIALOG(R)File 155:MEDLINE(R)

HIV-1 subtype O: epidemiology, pathogenesis, diagnosis, and perspectives of the evolution of HIV.

Gurtler LG; Zekeng L; Tsague JM; van Brunn A; Afane Ze E; Eberle J; Kaptue L

Max von Pettenkofer Institute for Hygiene and Medical Microbiology, University of Munich, Federal Republic of Germany.

Archives of virology. Supplementum (AUSTRIA) 1996, 11 p195-202, SSN 0939-1983 Journal Code: BLI

Languages: ENGLISH

Document type: Journal Article; Review; Review, Tutorial

Record type: Completed

HIV-1 subtype O is a new HIV variant originating in the West-Central African region, with highest prevalences in countries such as Cameroon, Equatorial Guinea and Gabon. Detection of antibodies to HIV-1 subtype O can pose problems in unmodified ELISA tests, and confirmation of anti-HIV-1 subtype O in immunoblot may give false negative results in some specimens. Nucleic acid-based assays designed for HIV-1 detection do not amplify or detect sequences from HIV-1 subtype O. In their env sequences, HIV-1 subtype O strains show a higher heterogeneity than the classical HIV-1 subtypes, leading to the conclusion that HIV-1 subtype O has been introduced into the human population only recently. Further, unidentified subtypes are also likely to exist. (15 Refs.)

Record Date Created: 19960927

17770

DIALOG(R)File 155:MEDLINE(R)

Reactivity of five anti-HIV-1 subtype O specimens with six different anti-HIV screening ELISAs and three immunoblots.

Gurtler LG; Zekeng L; Simon F; Eberle J; Tsague JM; Kaptue L; Brust S;

Max von Pettenkofer Institute, University of Munich, Germany,

Journal of virological methods (NETHERLANDS) Feb 1995, 51 (2-3) p177-83, ISSN 0166-0934 Journal Code: HQR

Languages: ENGLISH

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Document type: Journal Article

Record type: Completed

antigen/sandwich ELISA). One specimen was not detected by one peptide based ELISA. One specimen was only recognized by two ELISAs and should be completely, but showed some staining with the enzymes of HIV, and would commercially detected two of the specimens with a major binding of gp160 considered as a marker sample for the weakness of currently used ELISAs Five anti-subtype O specimens were tested by anti-HIV-1/2 screening and and a very weak staining of p24, and would most probably be interpreted as was also missed by most of the ELISAs, had very faint staining of the gp160 most probably be interpreted as indeterminate. The fifth specimen, which confirmatory assays. They can be divided into three specimens, reactive with all ELISAs, independent of the nature of the antigen (recombinant negative. Adaption of currently available tests to anti-subtype O is needed Another two specimens lacked reactivity with glycoproteins almost proteins or peptides) and test configuration (indirect ELISA or double with anti-subtype O. Three different immunoblot assays available and other viral bands, especially the integrase and reverse transcriptase. for the future reliability of anti-HIV diagnostic reagents.

Record Date Created: 19950606

DIALOG(R)File 155:MEDLINE(R)

Variability of human immunodeficiency virus type 1 group O strains solated from Cameroonian patients living in France.

Loussert-Ajaka I; Chaix ML; Korber B; Letourneur F; Gomas E; Allen E; Ly

ID; Brun-Vezinet F; Simon F; Saragosti S

Journal of virology (UNITED STATES) Sep 1995, 69 (9) p5640-9, ISSN Laboratoire de Virologie, Hopital Bichat-Claude Bernard, Paris, France. 0022-538X Journal Code: KCV

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

p24Gag and the Env C2V3 region were obtained from seven patients who were Human immunodeficiency virus type 1 (HIV-1) nucleotide sequences encoding Cameroonian HIV isolates of group O (HIV-1 outlier) than to group M (HIV-1 subtypes (A to H). Genetic distances between sequences from available group HIV enzyme-linked immunosorbent assay detection kits and having atypical Western blot (immunoblot) reactivity. Sequence analyses showed that all of large number of group M strains has to date revealed eight distinct genetic major). All seven patients had Cameroonian origins but were living in France at the time the blood samples were taken. Characterization of a selected on the basis of having paradoxical seronegativity on a subset of hese strains were more closely related to the recently described

sequence comparisons, showing that the group O viruses are genetically very diverse. Analysis of sequences from these seven new viral strains, combined they may have a comparable (or greater) age than the M group sequences, O isolates were generally comparable to those observed in M intersubtype sequences. The level of diversity among group O sequences suggests that discernable phylogenetic clustering patterns among the 10 patients' viral although for unknown reasons, the latter group dispersed first and is the with the three previously characterized group O strains, revealed few dominant lineage in the pandemic.

Record Date Created: 19950914

DIALOG(R)File 155:MEDLINE(R)

Isolation and envelope sequence of a highly divergent HIV-1 isolate: definition of a new HIV-1 group.

Charneau P; Borman AM; Quillent C; Guetard D; Chamaret S; Cohen J; Remy G Montagnier L; Clavel F

CNRS URA 1157, Departement Sida et Retrovirus, Institut Pasteur, Paris,

Virology (UNITED STATES) Nov 15 1994, 205 (1) p247-53, ISSN 0042-6822 Journal Code: XEA France.

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

HIV-1MVP5180, until now designated HIV-1 subtype O. Phylogenetic analysis solate from a French woman with AIDS. This virus, HIV-1VAU, is closely related to the recently described Cameroonian viral isolates HIV-1ANT70 and group). The HIV-1 group O is currently emerging in western central Africa more conventional HIV-1 subtypes. Therefore, these three viruses could be included in a new viral group, HIV-1 group O (outgroup), distinct from the We report here the isolation and envelope sequence of a divergent HIV-1 their mutual divergence is similar to what has been reported between the cluster of other HIV-1 isolates, which we will refer to as group M (Major reveals that the three viruses are equidistant from one another and that but its spread in Europe has already started.

Record Date Created: 19941206

07nov01 15:25:09 User208669 Session D1928.2

\$6.33 1.977 DialUnits File155

\$0.00 49 Type(s) in Format 6

\$2.00 10 Type(s) in Format 7

\$2.00 59 Types

\$8.33 Estimated cost File 155

**\$0.50 TYMNET** 

\$8.83 Estimated cost this search

\$9.09 Estimated total session cost 2.050 DialUnits

Logoff: level 01.10.01 D 15:25:09